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## **CLAIMS**

We claim:

- 1. A computer implemented method to search a heterogenous compound database for molecules which are likely to have the same biological activity as a known query molecule comprising the following steps:
  - a) defining fragments of a query molecule and a database molecules according to a defined set of rules;
  - b) generating shape descriptors for the query molecule and database molecule fragments; and
  - c) using the shape descriptors, identifying the database molecule which has a shape similar to the query molecule.
- 2. A computer implemented method to search a heterogenous compound database for molecules which are likely to have the same biological activity as a known query molecule comprising the following steps:
  - a) fragmenting a query molecule according to a defined set of rules;
  - b) topomerically aligning the query molecule fragments to generate a topomeric conformation;
  - c) generating the interaction energies between a probe and the atoms in the topomerically aligned query fragments at all intersection points in a three dimensional grid surrounding the aligned query fragments;
  - d) fragmenting a database compound according to a defined set of rules;

R. CRAMER, JILEK, LIU, GUESSREGEN, WENDT, AND K CRAMER

5

- e) topomerically aligning the database molecule fragments to generate a topomeric conformation;
- f) generating the interaction energies between a probe and the atoms in the topomerically aligned database fragments at all intersection points in a three dimensional grid surrounding the aligned query fragments;
- g) determining the similarity between query and database fragments by the root sum square differences in the field values; and
- h) identify the molecule in the database most similar to the query molecule as that molecule having the smallest field value difference in its fragments.